



Sequence	Length	Code
VLAEAMSQV (SEQ ID NO:70)	9	A
ILKEPVHGKV (SEQ ID NO:71)	9	B
TLNFPISPI (SEQ ID NO:72)	9	C
SLLNATDIAV (SEQ ID NO:73)	10	D
QMAVFIHNFK (SEQ ID NO:74)	10	E
VTVYYGVWPWK (SEQ ID NO:75)	11	F
FPVRPQVPL (SEQ ID NO:76)	9	G
YPLASLRSLF (SEQ ID NO:77)	10	H
VIYQYMDDLY (SEQ ID NO:78)	10	I
IYQEPFKNL (SEQ ID NO:79)	9	J
IWGCSGKLI (SEQ ID NO:80)	9	K

200
↓

202

AA	C+1 ranking	N-1 ranking
K	2.20	0.64
C	2.00	1.00
N	2.00	0.00
G	1.80	1.33
T	1.50	0.00
A	1.33	1.21
F	1.33	1.00
S	1.33	0.00
W	1.20	0.00
Q	1.20	0.00
R	1.17	1.57
M	1.00	0.00
Y	1.00	0.75
I	0.86	0.50
L	0.75	2.20
V	0.00	1.19
D	0.00	0.00
H	0.00	0.00
E	0.00	0.00
P	0.00	0.00

204

Motif Specification

XXXX(FY)XX(LIMV) (SEQ ID NOS:7-14)
 XXXX(FY)XXX(LIMV) (SEQ ID NOS:15-22)
 XXXXNXXX(LIMV) (SEQ ID NOS:27-30)
 XXXXNXXXX(LIMV) (SEQ ID NOS:341-344)
 X(LM)XXXXXXV (SEQ ID NOS:31-32) 206
 X(LM)XXXXXXXXV (SEQ ID NOS:33-34)
 X(LMVT)XXXXXX(KRY) (SEQ ID NOS:345-356)
 X(LMVT)XXXXXXXX(KRY) (SEQ ID NOS:357-368)
 XPXXXXXX(LIMVF) (SEQ ID NOS:59-63)
 XPXXXXXXXX(LIMVF) (SEQ ID NOS:64-68)

FIGURE 11A

Junctional Analyzer run on Saturday, February 26, 2000 09:06:23 pm.

The following non-zero AA weights will be used.

AA	N-1 ranking	C+1 ranking
A	1.21	1.33
C	1.00	2.00
F	1.00	1.33
G	1.33	1.80
I	0.50	0.86
K	0.64	2.20
L	2.20	0.75
M	0.00	1.00
N	0.00	2.00
Q	0.00	1.20
R	1.57	1.17
S	0.00	1.33
T	0.00	1.50
V	1.19	0.00
W	0.00	1.20
Y	0.75	1.00

204

The following 10 motif specifications will be used to search for junctionals.

Count	Motif Specification
1	XXXX(FY)XX(LIMV) (SEQ ID NOS:7-14)
2	XXXX(FY)XXX(LIMV) (SEQ ID NOS:15-22)
3	XXXXNXXXX(LIMV) (SEQ ID NOS:27-30)
4	XXXXNXXXX(LIMV) (SEQ ID NOS:341-344)
5	X(LM)XXXXXXV (SEQ ID NOS:31-32)
6	X(LM)XXXXXXXXV (SEQ ID NOS:33-34)
7	X(LMVT)XXXXXX(KRY) (SEQ ID NOS:345-356)
8	X(LMVT)XXXXXXXX(KRY) (SEQ ID NOS:357-368)
9	XPXXXXXX(LIMVF) (SEQ ID NOS:59-63)
10	XPXXXXXXXX(LIMVF) (SEQ ID NOS:64-68)

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Code	Peptide	Length
A	VLAEAMSQV	(SEQ ID NO:70)
B	ILKEPVHGV	(SEQ ID NO:71)
C	TLNFPISPI	(SEQ ID NO:72)
D	SLLNATDIAV	(SEQ ID NO:73)
E	QMAVFIFHNFK	(SEQ ID NO:74)
F	VTVYYGVPVWK	(SEQ ID NO:75)
G	FPVRPQVPL	(SEQ ID NO:76)
H	YPLASLRSLF	(SEQ ID NO:77)
I	VIYQYMDDLY	(SEQ ID NO:78)
J	IYQEPFKNL	(SEQ ID NO:79)
K	IWGCSGKLI	(SEQ ID NO:80)

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MaxInsertions = 4 (208)

FIGURE 13A

EP-HIV-1090 (SEQ ID NO:81)

MGMQVQIQSLFLLLLWVPGSRGKLVGKLNWAGAAILKEPVHGVNAACPVSFEPKIPHYCAPA
KAKFVAATLKAAKAFAFPVRPQVPLGAALKTLCVTLGAAVLAEAMSQVKVYLAWVPAHKG
AAAAIFQSSMTKKTTLFCASDAKNIPYNPQSQGVVKHPVAGPIANVTYYGVPVWKAAAQMA
VFIHNFKNAAAAYPLASLRLFNLTFGWCFKLNRLQQLFINAKIQNFRVYYRKAATIKIGGQLKK
VPLQLPLKAMTNNNPPPIP

ATGGGAATGCAGGTGCAGATCCAGAGCCTGTTCTGCTCCTCCTGTGGGTGCCGGATCCAGA
GGAAAGCTGGTGGCAAACACTCAACTGGGCCGGAGCTGCAATCTGAAGGAGCCCGTCCACGG
GGTGAATGCCGCTGCCCTAAAGTCAGCTTCAACCATAAGATCCCCATTCAATTACTGTGC
ACCTGCCAAAGCTAAGTTGTGGCCGCTGGCACCTCAAGGCCGCTGCAAAAGCCTTCCAGT
GAGGCCAGGTGCCTCTGGCGCCGCTAAACACTCACACCAGTGCCTGACTCTGGGAGCCGC
TGCAGTGCTGGCAGAGGCCATGTCCAAGTGAAGGTGTATCTGGCITGGGTGCCGCCACAA
GGGGGCCGCTGCAGCCATCTTCAGTCTAGCATGACCAAGAAAACAACCTCTGTTCTGTGCCTC
CGACGCTAAGAACATCCCTTATAATCCACAGTCAGGGCGTGGTCAAGCATCCGTGCACGC
CGGACCTATTGCTAACGTGACCGTGACTATGGGTCCCAGTGTGAAAGAAAGCCGCTGCACA
GATGCCGTGTTATTACAATTCACAAAACGCCGCTGCATACCCCTGCCAGCCTGAGATC
CCTCTCAACCTGACATTGGCTGGTCTTAAGCTGAACCGGATCTGCAGCAACTGCTCTT
ATCAATGCTAAAATCCAGAACTTCCGCTACTATAGGAAGGCTGCAGTGAATGACTATCAAATT
GGCGGACAACCTGAAGAAAGTGCCTCTCCAGCTGCCCTCTCAAGGCAATGACCAACAATCC
CCCTATCCCAGTCTGA (SEQ ID NO:82)

HIV-CPT (SEQ ID NO:83)

MGMQVQIQSLFLLLLWVPGSRGIPHYCAPAKAAKIQNFRVYYRKAATIKIGGQLKKAKFVAAW
TLKAAAKVPLQLPPLKAIFQSSMTKKTLCVTLGAQMAVFIFHFKGAKVYLAWVPAHKNAIPYN
PQSQGVVKAILKEPVHGVGAAALTFGWCFKLNAVLAEAMSQVNRLQQLFINAAACPVSFEP
KVTYYGVPVWKAAHPVHAGPIANAAAAYPLASLRLFNAAATTLCASDAKNKLVGKLNWAN
AAAFPVRPQVPLNMNNPPPIP

ATGGGGATGCAGGTGCAGATCCAGAGCCTGTTCTGCTCCTCCTGTGGGTGCCGGATCCAGA
GGAATCCCCATTCACTACTGCGCCCTGCTAAGGCAGC AAAATCCAGAAACTCAGGGTGTAT
TACAGAAAGGCTGCAGTCACCATTAAATCGCGGACAACCTGAAGAAAGCCAAGTTGTGGC
CGCTTGGACACTCAAGGCCGCTGCAAAGGTCCCAGTGCAGCTCCCCCTCTGAAGGCCATCTT
CCAGAGCTCATGACTAAGAAACTGACCCACTGTGTGACACTCGGGGCCAGATGGCTGT
GTTCATCCATAATTAAAGGCGCCAAGGTCTACCTGGCTGGGTGCCGCACACAAGAACGC
CATTCCCTACAATCCACAGTCTCAAGGAGTGGTCAAAGCTATTCTGAAGGAGCCGTGCACGG
GGTGGCGCCGCTGCACTCACTTGGATGGTCTTAAACTGAACGCCGTGCTGGCTGAAGC
CATGAGCCAGGTCAATCGGATCCTGCAGCAACTGCTCTCATTACGCCGCTGCATGTCCTAA
GGTGTCTTCGAGCCAATCAAAGTGACCGTGATTACGGGGTCCCCGTGTGAAAGAAAGCCGC
TCATCCTGTCCACGCAGGCCAATGCCAACGCCGCTGCATATCCCTGCCCTCTGCCAG
CCTGTTAACGCCGCTGCAACAACCCCTTTGCGCCTCCAGCTAAGAATAAAACTGGGG
AAAGCTGAACCTGGCCAACGCAGCTGCATTCCCTGTGAGGCCACAGGTCCCCCTCAATATGAC
TAACAATCCCCCTATCCCAGTGTGA (SEQ ID NO:84)

FIGURE 18A

HIV-FT (SEQ ID NO:85)

MQVQIQSLLLLWVPGSRGKLVGKLNWAMASDFNLPPVAIFQSSMTKVTIKIGGQLKRILQQLLF
IMAVFIHNFKIPYNPQSQGVVTTLFCASDAKILKEPVHGVQMAVFIHNFKGAAVFIHNFKRCPKVSF
EPIKIQNFRVYRLLTFGWCFLQVPLRPMTYKMTNNPPIPVTVYYGVPVWKVLAEAMSQVIPHY
CAPAKLTPLCVTL

ATGCAGGTGCAGATCCAGAGCCTGTTCTGCTCCTCTGGGTGCCGGATCCAGAGGAAG
CTGGTGGGAAGCTGAACCTGGCCATGGCCAGCGATTCAACCTGCCCGTGGCCATCTTC
CAGAGCAGCATGACCAAGGTGACCATCAAGATCGGGGGCAGCTGAAGAGGATCCTGCAGCA
GCTGCTGTTCATCATGGCGTGTTCATCCACAACCTCAAGATCCCCTACAACCCCCAGAGCCA
GGGGGTGGTGACCACCCCTGTTCTGGCCAGCGATGCCAAGATCTGAAGGAGCCCGTGCACG
GGGTGCAGATGGCGTGTTCATCCACAACCTCAAGGGCGCCGCGTGTTCATCCACAACCTCA
AGAGGTGCCCAAGGTGAGCTCGAGCCCATCAAGATCCAGAACCTCAGGGTGTACTACAGG
CTGACCTTCGGGTGGTGTTCAAGCTGCAGGTGCCCTGAGGCCATGACCTACAAGATGACC
AACAAACCCCCCATCCCCGTGACCGTGTACTACGGGGTGCCCGTGTGGAAGGTGCTGGCGAG
GCCATGAGCCAGGTGATCCCCATCCACTACTGCGCCCCGCCAAGCTGACCCCCCTGCGTG
ACCCTG (SEQ ID NO:86)

FIGURE 18B

HIV-TC (SEQ ID NO:87)

MGMQVQIQSLFLLLLWVPGSRGYWQATWIPEWKAIFQSSMTKKVYLAWVPAHKNAACPVSFE
PIKHPVHAGPIANLTGWCFLNKMIIGGIGGFIKFRDYVDRFYKAAARILQQLFINTTLCASDAK
NQMVKHQAIISPRGAALKVGKLNWAGAAAIIYETYGDTWKAACQVPLRPMTYKGAAAVTLDVGDAY
NAAARYLKDQQLNTLNFPISPINMTNNPIPVNAPYNTPVFAIKAAAVPLQLPPLKAAIPYNPQSQ
GVVKALLQLTVWGIGAAILKEPVHGvnAAAFPISPIETVKVWKEATTLFKAAAVTIKIGGQLKKI
YQEFPKNLKAAAALVLAEAMSQVNVLGPTPVNIGAAAEVNITDSQYKAAAIPHYCAPAKAVIYQY
MDDLYKAAAQMAVFIFHNFKNAATYQIYQEPFKPYNEWTLELKAKIQNFRVYRKAFTPVRPQVPL
GAAAIWGCSGKLIKVMIVWQVDRNAAKAACWWAGIKAKFVAATLKAALTPLCVTLNAAM
ASDFNLPPVKSLLNATDIAVNVTVYYGVPVWKAAAIIIRILQQLKRAMASDFNLNAAYPLASL
RSLF

ATGGGGATGCAGGTGCAGATCCAGAGCCTGTTCTGCTCCTCTGGGTGCCGGATCTAGA
GGATACTGGCAAGCTACTTGGATTCCAGAACAGAAAGCTATCTTCAATCCTCAATGACGAAG
AAGGTATACCTGGCATGGGTCCCAGCACACAAGAACGCCGCTGCCAAAGGTGTCTTGA
CCCCATTAAACACCCAGTCACGCAGGGCCAATAGCGAATTGACATTGGGTGGTGTCTCAA
CTAAACAAAATGATCGCGGCATTGGAGGCTTATCAAGTTAGAGATTACGTGGACCGATT
TATAAAAGCCGCTGCCGTATACTCCAGCAGCTACTATTCACAAACACCCTCTCTGCGCTT
CAGACGCTAAGAACCAATGGTACACCAAGCCATAAGCCCTAGAGGAGCCAAGCTCGTAGGG
AAATTAAATTGGCGGGTGCAGCAGCAATCTACGAGACTTACGGCGATACTGGAAAGCAGC
CCAGGTTCCGTTACGCCAATGACCTATAAAGGCGCAGCAGCTAACAGTTAGATGTAGG
AGACGCTTACAACGCTGCCGCAAGATACCTAAAGATCAGCAGTTACTCAACACACTAAATT
CCCAATTAGCCGATAAACATGACAAATAACCCACCAATTCCGTCATGCTCCCTACAACAC
TCCAGTATTGCAATCAAAGCCGCTGCTGCCCCCTGCAGCTCCCTCTGAAAGCTGCGAT
ACCTTACAACCCACAGAGCCAAGGTGTGTCAAAGCACTGCTCAGCTAACAGTTGGGAAT
TGGTGTGCAATTCTAAAGAGCCAGTTACGGGTTAACGCCGCCCTCCCAATCAGTCC
TATTGAGACTGTGAAAGTATGAAAGAACCCACAACACTTTTAAGGCAGCCAGTTA
CAATTAAAATAGGGGCCAACCTAAGAAAATATACCAGGAACCTTCAAGAATCTCAAAGCC
GCTGCAGTGTGCGCCGAGGCTATGTCACAGGTGAATTGGTGGACCAACACCGTAAACATC
GGAGCCGAGCCGAAGTGAACATAGTCACCGACTCACAGTACAAAGCCGCTGCAATACCCAT
ACATTATTGTGCTCCGCAAAGGCCGTATCAATATATGGACGACCTGTATAAGGCCGC
CGCGCAGATGGCAGTCTTATCCACAACCTTAAAGCAGCTACTTATCAGATCTACAGGA
ACCATTCAAACCGTACAATGAGTGGACCTTGGAACTAAAGGCCAAATTCAGAACCTCAGGG
TATATTATAGAAAAGCATTCCAGTGAGGCCCAAGGTGCTCTGGGTGCCGCAGCAATATGGG
GATGTTCTGGAAAAGTGTCAAGGTGATGATTGTATGGCAAGTGGACAGAAAATGCAGCTAAG
GCAGCCTGTTGGTGGCAGGTATAAAAGCAAAGTTCGTGGCAGCATGGACGCTTAAAGCAGC
CGCAAAACTCACTCCTCTGCGTGACACTTAATGCAGCCATGCCCTCTGATTCAACCTCCC
CCTGTAATCCCTGCTTAATGCGACAGATATCGCAGTCACGTAACAGTATATTATGGCGTG
CCAGTCTGGAAAAAGCCGCCGCGCCATAATTGGATACTGCAGCAGCTGAAAAGAGCTAT
GGCGAGTGACTCAACCTGAATGCGGCCGCTACCCCTGGCATCGTTAAGGTCACTATTTG
A (SEQ ID NO:88)

FIGURE 18C

HCV.1 (SEQ ID NO:89)

MGMQVQIQSLFLLLWVPGSRGLFNILGGWVDLMGYIPLVYLVAYQATVILAGYGAGVRLIVFP
DLGVHMWNFISIGIYLLPREGPRLYLVTTRADVVLVGGVLAALLFLLADAFLLADARVWMNRL
IAFACTCGSSDLYLSAFLSHSYGVAGALVAFKLPGCSFSIFKTSERSQPRLIFCHSKKFWAKHMW
NFIPFYGKAIRMYVGGVEHRQLFTSPRRRLGVRATRKVGIYLLPNRAKFVAAWTLKAAA*

GAATTGCCGCCACCATGCAGGTGCAGATCCAGAGCCTGTTCTGCTCCTCCTGGGGTCCCCGGATCCAGAGGACTGC
TGGTCACATCCTGGGGGGTGGATCTGATGGGGTACATCCCCCTGGTGTACCTGGTGGCTACCAGGCCACCGT
GATCCTGGCCGGTAACGGGCCCCGGTGGAGCTGATCGTGTCCCCGATCTGGGGTGCACATGTGAACCTTCATCAGC
GGGATCTACCTGCTGCCAGGAGAGGACCTAGACTGTACCTGGTGACTAGACACGCTGATGTGGTGTGGGGAGGAG
TGCTGGCTGCTCTGCTGTTCTGCTGGCTGATGCTTCTGCTGCTGGCTGATGCTAGAGTGTGGATGAACAGACT
GATCGCTTCGCTTGATCATGTGGAAGCTCCGATCTGTATCTGAGCGCTTCAGCCTGCACAGCTACGGAGTGGCTGGA
GCTCTGGTGGCTTTAAGCTGCCGGATGTAGCTTAGCATCTTAAGACCAGCGAAAGAACGCCAGCCTAGACTGATCT
TTTGTCAACAGCAAGAAGAAGTGGGCTAACACATGTGAATTATCCCTTCTATGAAAGGCTATCAGAATGTA
TGTGGGAGGAGTGGAACACAGACAGCTGTTACATTAGCCCTAGAAGGAGACTGGGAGTGAGAGCTACAAGAAAGGTG
GGAATCTATCTGCTGCCAAATAGATGAAAGCTGGG* (SEQ ID NO:90)

HCV.2 (SEQ ID NO:91)

MGMQVQIQSLFLLLWVPGSRGDLMGYIPLVAKFVAAWTLKAAALLFLLADALIFCHSKKKQLF
TFSPRRLVTRHADVYLLPREGPRLCGSSDLYHMWNFISIGIFWAKHMWNFAKFVAAWTLKAA
AILAGYGAGVYLVAYQATVGVAGALVAFKIPFYGKAIRMYVGGVEHRVLVGGVLAFLLLADA
RVLPGCSFSIFAKFVAAWTLKAAAKTSERSQPRRLGVRATRKRLIVFPDLGVWMNRLIAFALSafs
LHSYLLFNILGGWVVGICYLLPNR*

GAATTGCCGCCACCATGGGAATGCAGGTGCAGATCCAGAGCCTGTTCTGCTCCTCCTGTGG
GTGCCGGATCCAGAGGAGATCTGATGGGATATATCCTCTGGTGGCTAACGTTGTGGCTGCT
TGGACACTGAAGGCTGCTGCTGCTGCTGCTGCTGGCTGATGCTCTGATCTCTGTCACA
GCAAGAAGAACGAGCTGTTACATTAGCCCAGAAGATATCTGGTACAAGAACACGCTGAT
GTGTATCTGCTGCCAGACGGGACCTAGACTGTGATCTGGAAGCTCCGATCTGTATCAC
ATGTGGAACCTTATCAGCGGAATCTTGGGCTAACGACATGTGGAATTCTCATCCTGGCTGGA
TATGGAGCTGGAGTGTATCTGGTGGCTTATCAGGCTACAGTGGGAGTGGCTGGAGCTCTGGT
GCTTCAAGATCCCATTCTATGAAAGGCTATCAGAATGTATGTGGAGGAGTGGAACACAG
AGTGCTGGTGGGAGGAGTGTGGCTGCTGCTGCTGGCTGATGCTAGAGTGTGCCAGG
ATGTAGCTTAGCATCTCAAGACTCCGAACGCTCCCAGCCTAGAAGACTGGGAGTGAGAGC
TACAAGGAAGAGACTGATCGTGTCTCAGATCTGGGAGTGTGGATGAATAGACTGATCGCTT
CGCTCTGAGCGCTTCAAGCTGCAAGCTATCTGCTGTTCAACATCCTGGGAGGATGGGTGGT
GGGAATCTATCTGCTGCCAAACAGATGAAAGCTT (SEQ ID NO:92)

HCV.3s1 (SEQ ID NO:93)

MGMQVQIQSLFLLLWVPGSRGYLVAYQATVAKFVAAWTLKAAALLFLLADALIFCHSKKKYL
VTRHADVLGFAYMSKCTCGSSDLYHMWNFISIGIFWAKHMWNF*

GAATTGCCGCCACCATGGGAATGCAGGTGCAGATCCAAAGCCTGTTCTGCTCCTCCTGTGG
GTGCCGGATCCAGAGGATACCTCGTCGCCTACCAGGCCACTGTGGCTAACATTGTCAGCC
TGGACACTGAAGGCTGCACTGCTGCTCTGCTGCTGGCCATGCACTCATCTCTGCCATT
CCAAGAAAAAGTATCTGGTCAACAGACATGCTGACGTGCTGGGTTGGCGCTACATGAGC
AAGTGCACCTGTGGCAGCTCCGACCTGTATCACATGTGGAACCTTATTCTGGAATCTTGGG
CCAAGCACATGTGGAATTCTGAAAGCTT (SEQ ID NO:94)

FIGURE 18D

HCV.3s2 (SEQ ID NO:95)

MGMQVQIQSLFLLLLWVPGSRGVLVGGVLAAAKFVAAWTLKAAAFLLADARVLSAFSLHSYIL
AGYGAGVWMNRLIAFAIPFYGKAIVAGALVAFKVGIYLLPNR*

GAATTGCCGCCACCATGGGAATGCAGGTGCAGATCAAAGCCTGTTCTGCTCCTCCTGTGG
GTGCCCAGATCCAGAGGAGTCCTGGTGGCGCGCTCTGGCGCTGCTAAGTTGTCGCTGCT
TGGACACTGAAGGCAGCCGCTTCTGCTCCTGGCAGACGCCAGGGTGTCTGCCTTCAGC
CTCCACTCCTACATCCTCGCAGGTATGGCGAGCGTGTGGATGAATCGGCTGATGCCCTT
GCCATTCCATTCTATGGAAAGCCATTGTGGCTGGCGCCCTGGTGGCATTCAAGGTCGGGATC
TACCTCCTGCCTAACCGCTGAAAGCTT (SEQ ID NO:96)

HCV.3s2(-3) (SEQ ID NO:97)

MGMQVQIQSLFLLLLWVPGSRGVLVGGVLAAAKFVAAWTLKAAAFLLADARVLSAFSLHSYIL
AGYGAGVWMNRLIAFA*

GAATTGCCGCCACCATGGGAATGCAGGTGCAGATCAAAGCCTGTTCTGCTCCTCCTGTGG
GTGCCCAGATCCAGAGGAGTCCTGGTGGCGCGCTCTGGCGCTGCTAAGTTGTCGCTGCT
TGGACACTGAAGGCAGCCGCTTCTGCTCCTGGCAGACGCCAGGGTGTCTGCCTTCAGC
CTCCACTCCTACATCCTCGCAGGTATGGCGAGCGTGTGGATGAATCGGCTGATGCCCTT
GCCCTGAGGATCC (SEQ ID NO:98)

HCV.3s3 (SEQ ID NO:99)

MGMQVQIQSLFLLLLWVPGSRDLMGYIPLVAKFVAAWTLKAAARLGVRATRKLLFNILGGWV
RMYVGGVEHRRLIVFPDLGVGVAGALVAFKLPGSFSIFKTSERSQPRQLFTSPRRYLLPREGPRL

GAATTGCCGCCACCATGGGAATGCAGGTGCAGATCAAAGCCTGTTCTGCTCCTCCTGTGG
GTGCCCAGATCCAGAGGAGACCTGATGGGCTACATCCCTCTCGTGGCCAAGTTGTCGAGCT
TGGACCTGAAAGCCGCTGCCAGACTGGGAGTGCGCGCTACACGAAACTCCTGTTAACATC
CTGGGAGGGTGGGTGCGGATGTACGTCGGAGGGCGTCAGCACAGAAGGCTATTGCTTCC
AGATCTGGCGTGGCGTCGAGCGACTCGTGGCTCAAACTGCCAGGGTGCAGCTTCAG
CATTTCAAGACCTCCGAACGCTCCCAACCCAGACAGCTTCACTTCTCCTCGGAGGTAT
CTGCTGCCAGACCGGACCCAGGCTGAAAGCTT (SEQ ID NO:100)

HCV.PC3 (SEQ ID NO:101)

MGMQVQIQSLFLLLLWVPGSRGLLFNILGGWVKAKFVAAWTLKAAALADGGCSGGAYRLIVFPD
LGVKFWAKHMWNFIGVAGALVAFKKQLFTSPRR*

GAATTGCCGCCACCATGGGAATGCAGGTGCAGATCAAAGCCTGTTCTGCTCCTCCTGTGG
GTGCCCAGATCCAGAGGAGTCCTGCTTCAACATCCTGGCGGATGGGTGAAGGCCAAGTCGCTG
GCTGCCTGGACCTGAAGGCTGCCGCTCTGGCGACGGGGATGCAGCGGCGAGCTACAG
GCTCATTGCTTCCCAGTCGGAGTCAATTGGCAAAGCACATGTGGAATTTCATCGG
GGTGGCCGGAGCCCTGGTCGTTAAAAGCAGCTTCACCTCTCCCAAGACGGTGAGG
TACC (SEQ ID NO:102)

FIGURE 18E

HCV.PC4 (SEQ ID NO:103)

MGMQVQIQSLFLLLLWVPGSRGRLGVRATRKAKFVAAWTLKAAAKTSERSQPRNLPGCSFSIFNDLMGYIPLVKYLLPRRGPRLNTLCGFADLMGYRMYVGGVEHR*

GAATTGCCGCCACCATGGGAATGCAGGTGCAGATCAAAGCCTGTTCTGCTCCTCCTGTGG
GTGCCCGGATCCAGAGGAAGGCTGGCGTGAGAGGCCACCCGAAGAAGGCCAAGTTCGTGGC
TGCCTGGACCCCTGAAGGCTGCCGCTAAAACAAGCGAGCGCTCCCAGCCCAGGAACCTGCCTG
GATGCTCTTCAGCATCTTAATGACCTCATGGGTACATTCCACTGGTGAAGTATCTGCTCCC
CAGACGGGGCCCTCGCCTGAACACTCTGTGGATTGCTGATCTGATGGGTACAGGATGTA
TGTCCGGCGAGTCGAACACAGATGAGGTACC (SEQ ID NO:104)

HCV.2431(1P) (SEQ ID NO:105)

MGMQVQIQSLFLLLLWVPGSRGVLVGGVLAAAFLLADARVLSAFSLHSYILAGYGAGVWMNRL
IAFAGAAAARLGVRATRKAAAKTSERSQPRNLPGCSFSIFNDLMGYIPLVKYLLPRRGPRLNTLCG
FADLMGYRMYVGGVEHRKLLFNILGGWVKAALADGGCSGGAYRLIVFPDLGVKFWAKHMWN
FIGVAGALVAFKKQLFTSPRRNGYLVAYQATVAAALLFLLADALIFCHSKKKYLVTRHADVLG
FGAYMSKCTCGSSDLYHMWNFISGIFWAKHMWNKAAAACKVAAWTLKAAA

GAATTGCCGCCACCATGGGAATGCAGGTGCAGATCAAAGCCTGTTCTGCTCCTCCTGTGG
GTGCCCGGCTCCAGAGGAGTCCTGGTGGCGCGTCTGGCAGCCGCTTCTGCTCCTGGCA
GACGCCAGGGTGCCTGCTGCCTCAGCCTCACTCCTACATCCTCGCAGGGTATGGCGCAGGC
GTGTGGATGAATCGGCTGATGCCCTTGGCGCTGCCGCAAGGCTGGCGTGAGAGGCCACC
CGGAAGAAGGCTGCCGCTAAAACAAGCGAGCGCTCCCAGCCAGGAACCTGCTGGATGCTC
TTTCAGCATCTTAATGACCTCATGGGTACATTCCACTGGTGAAGTATCTGCTCCCCAGACGG
GGCCCTCGCCTGAACACTCTGTGGATTGCTGATCTGATGGGTACAGGATGTATGTCGGC
GGAGTCGAACACAGAAAATGCTCTCAACATCCTGGCGGATGGGTGAAGGCTGCCGCTCT
GGCCGACGGGGATGCAGCGCGGAGCTTACAGGCTCATTGTCTTCCGATCTCGGAGTC
ATTTGGCAAAGCACATGTGAATTTCATGGGTTGGCGGAGCCCTGGTCGCTTTAAAAAA
GCAGCTCTCACCTCTCCCCAAGACGGAACGGATACCTCGCCTACCAAGGCCACTGTGGC
TGCAGCTCTGCTCTCCTGCTGCCGATGCACTCATCTCTGCCATTCCAAGAAAAGTAT
CTGGTCACCAGACATGCTGACGTGCTGGGTTGGCGCCTACATGAGCAAGTGCACCTGTGGC
AGCTCCGACCTGTATCACATGTGAATTTCATGGGAACTTATTTCTGGATCTTGGCCAAGC
AATTTAAGGCCGAGCTAAATTGTCGGCAGCCTGGACACTGAAAGCAGCTGCATGAGG
ATCC (SEQ ID NO:106)

FIGURE 18F

HCV.4312(1P) (SEQ ID NO:107)

MGMQVQIQSLFLLLLWVPGSRGRLGVRATRKAAAKTSERSQPRNLPGCSFSIFNDLMGYIPLVK
YLLP RRGPRLNTLCGFADLMGYRMYVGGVEHRKLLFNILGGWVKAALADGGCSGGAYRLIVFP
DLGVKF WAKHMWNFIGVAGALVAFKKQLFTSPRRNGYL VAYQATVAALLFLLLADALIFCHS
KKKYLVTRHADVLGFGAYMSKCTCGSSDLYHMWNFISGF WAKHMWNFKKAALVGGVLAA
AFLLLADARVLSAFSLHSYILAGYGAGVWMNRLIAFANAAAKFVAAWTLKAAA*

GAATTCCGCCACCATGGGAATGCAGGTGCAGATCAAAGCCTTTCTGCTCCTCTGTGG
GTGCCCGCTCCAGAGGAAGGCTGGCGT GAGAGCCACCCGGAAAGAAGGCTGCCGTAAAAC
AAGCGAGCGCTCCAGCCCAGGAACCTGCCTGGATGCTCTTCAGCATTTAATGACCTCAT
GGGTACATTCACTGGTAAGTATCTGCTCCCCAGACGGGGCCCTGCCCTGAACACTCTG
TGGATTGCTGATCTGATGGGTACAGGATGTATGTCGGCGGAGTCGAACACAGAAAATGCT
CTTCAACATCCTGGCGGATGGGTGAAGGCTGCCCTGCCGACGGGGATGCAGCGCG
GAGCTTACAGGCTCATTGTCTTCCGATCTGGAGTCAAATTGGCAAAGCACATGTGGA
ATTTCATCGGGTGGCCGGAGCCCTGGTCGCTTTAAAAAGCAGCTCTCACCTCTCCCCAA
GACGGAACGGATACCTCGCCTACCAGGCCACTGTGGCTGCAGCTCTGCTCTGCTCC
TGGCCGATGCACTCATCTCTGCCATTCCAAGAAAAGTATCTGGT CACCAGACATGCTGACG
TGCTGGGTTTGGCGCTACATGAGCAAGTGCACCTGTGGCAGCTCCGACCTGTATCACATGT
GGAACTTATTCGGAATCTTGGCAAGCACATGTGGATTITAAGAAAGCCGCTGCAG
TCCTGGTGGCGCGCTGGCAGCCCTTCTGCTCTGGCAGACGCCAGGGTGTCTG
CCTTCAGCCTCCACTCCTACATCCTCGCAGGTATGGCGCAGGCGTGTGGATGAATCGGCTGA
TCGCCTTGCCAATGCTGCAGCTAAATTGTGGCAGCCTGGACACTGAAAGCAGCTGCATGAG
GATCC · (SEQ ID NO:108)

AOSL.K (SEQ ID NO:109)

MGMQVQIQSLFLLLLWVPGSRGHTLWKAGILYKAKFVAAWTLKAAAFLPSDFFPSVKFLSLGIH
LYMDDVVLGVGLSRYVARLFLTRILTISTLPETTVRRQAFTFSPTYKWLSSLVPFV

ATGGGAATGCAGGTGCAGATCCAGAGCCTTTCTGCTCCTCTGTGGGTGCCCGGGTCCAGA
GGACACACCCCTGTGGAGGCCGAATCCTGTATAAGGCCAAGTTCGTGGCTGCCTGGACCCCTG
AAGGCTGCCGTTCTGCCTAGCGATTCTTCTAGCGTGAAGTCTCTGCTGTCCCTGGAA
TCCACCTGTATATGGATGACGTGGTGTGGAGTGGACTGTCCAGGTACGTGGCTAGGCTGT
TCCTGCTGACCAGAACCTGACCATCTCCACCCCTGCCAGAGACCACCGTGGT GAGGAGGCAGG
CCTCACCTTAGCCCTACCTATAAGTGGCTGAGCCTGCTGGTGCCTTGTGA (SEQ ID NO:110)

HBV.1 (SEQ ID NO:111)

MGMQVQIQSLFLLLLWVPGSRGHTLWKAGILYKAKFVAAWTLKAAAFLPSDFFPSVKFLSLGIHL
YMDVVVLGVGLSRYVARLFLTRILTISTLPETTVRRQAFTFSPTYKWLSSLVPFVIPSSWAFTP
ARVTGGVFVKVGNFTGLYLPSPDTFFPSVTLWKAGILYKNVSIPWTHKLVVDQSFRSAICSVVRRAL
MPLYACI

ATGGGAATGCAGGTGCAGATCCAGAGCCTTTCTGCTCCTCTGTGGGTGCCCGGGTCCAGA
GGACACACCCCTGTGGAGGCCGAATCCTGTATAAGGCCAAGTTCGTGGCTGCCTGGACCCCTG
AAGGCTGCCGTTCTGCCTAGCGATTCTTCTAGCGTGTCTGCTGTCCCTGGAAATCC
ACCTGTATATGGATGACGTGGTGTGGAGTGGACTGTCCAGGTACGTGGCTAGGCTGTCC
TGCTGACCAAGAACCTGACCATCTCCACCCCTGCCAGAGACCACCGTGGT GAGGAGGCAGGCCT
TCACCTTAGCCCTACCTATAAGTGGCTGAGCCTGCTGGTGCCTTGTGATCCCTATCCCTAG
CTCCTGGGCTTACCCCAAGCCAGGGTGACCGGAGGAGTGTAAAGGTGGAAACTTCACCGG
CCTGTATCTGCCAGCGATTCTTCTAGCGTGA CCCTGTGGAAAGCCGGATCTGTACAA
GAATGTGTCCATCCCTGGACCCACAAGCTGGTGGACTTTCCAGTTCAGCAGATCCGC
TATCTGCTCCGTGGT GAGGAGAGCTCTGATGCCACTGTATGCCGTATCTGA (SEQ ID NO:112)

FIGURE 18G

HBV.2 (SEQ ID NO:113)

MGMQVIQSLFLLLWVPGSRGHTLWKAGILYKAKFVAAWTLKAAFLPSDFPSVNFLSLGIHLYMDDVVLGVGLSRYVARLFLLTRILTISTLPETTVVRRQAFTFSPYKGAAAWSLLVPFVNIPIPSSWAFKTPARVTGGVFVKVGNFTGLYNLPSDFPSVKTLWKAGILYKNVSIPWTHKGAALVVDFSQFSRNSAICSVVRRALMPYACI

ATGGGAATGCAGGTGCAGATCCAGAGCCTGTTCTGCTCCTCCTGTGGGTGCCCGGATCCAGA
GGACACACCCCTGTGGAAGGCCCGAATCCTGTATAAGGCCAAGTTCGTGGCTGCCTGGACCTG
AAGGCTGCCGCTTCTGCTAGCGATTCTTCCTAGCGTGAACCTCCTGCTGTCCCTGGAA
TCCACCTGTATATGGATGACGTGGTCTGGGAGTGGGACTGTCCAGGTACGTGGTAGGCTGT
TCCTGCTGACCAGAACATCCTGACCATCTCACCCCTGCCAGAGACCACCGTGGTAGGAGGCAGG
CTTCACCTTTAGCCCTACCTATAAGGGAGCCGCTGCCTGGCTGAGCCTGCTGGTGCCTTGT
GAATATCCCTATCCCTAGCTCCTGGGCTTCAAGACCCCAGCCAGGGTACCGGAGGAGTGT
TAAGGTGGAAACCTCACCGGCCTGTATAACCTGCCAGCGATTCTTCCTAGCGTGAAGAC
CCTGTGGAAGGCCCGAATCCTGTACAAGAAATGTGTCATCCCTGGACCCACAAGGGAGCCG
CTCTGGTGGACTTTCCCAGTCAGCAGAAATTCCGCTATCTGCTCCGTGGTGAGGAGAG
CTCTGATGCCACTGTATGCCGTATCTGA (SEQ ID NO:114)

PfCTL.1 (SEQ ID NO:115)

MQVQIQSLFLLLWVPGSRGILSVSSFLFVNAAAQTNFKSLRNLPSENERGYKAAALLACAGLAY
KKAACAKFVAAWTLKAAAKAFMKAVCVEVNAASFLFVEALFNATPYAGEPAPFKAAKYKLA
TSVLKAGVSENIFLKNAAYFILVNLLIKAGLLGVVSTV

ATGGGAATGCAGGTGCAGATCCAGAGCCTGTTCTGCTCCTCCTGTGGGTGCCCGGATCCAGA
GGAATCCTGAGCGTGTCCCTTCTGTTGTCAACGCCGCTGCACAGACCAATTCAAGAGC
CTCCTGAGGAACCTCCCTCCGAGAACGAAAGAGGCTACAAAGCCGCTGCACTGCTGCCCTGC
GCTGGACTGCCCTATAAGAAAGCCGCTGCAGCCAAGTTCGTGGCCGCTTGGACACTGAAGGC
CGCTGCAAAGCCTTATGAAGGCTGTCTGTGGAGGTCAATGCCGCTGCATCTTCTGT
GTGGAGGCCCTTTAACGCTACTCCTTACGCAGGGGACCAGCCCCCTCAAGGCCGCTGCA
AAATATAAGCTGCAACCAGCGTGTGAAGGCTGGCGTCCGAGAATATTTCTGAAAAAC
GCCGCTGCATACTCATCTGGTGAATCTGCTATTAGGCCGACTCCTGGGGTGGTCTCT
ACAGTGTGA (SEQ ID NO:116)

PfCTL.2 (SEQ ID NO:117)

MQVQIQSLFLLLWVPGSRGFVEALFQEYNAAAKYLVIVFLINALACAGLAYKKFYFILVNLLKA
ALFFIIFNKNAAKFVAAWTLKAAAKFILVNLLIFHNFQDEENIGIYKLPYGRNLKAAAVLLGGV
GLVLFNLIFFDLFLVKAVLAGLLGVV

ATGGGAATGCAGGTGCAGATCCAGAGCCTGTTCTGCTCCTCCTGTGGGTGCCCGGATCCAGA
GGATTCTGGAGGCCCTGTTCAAGGAATACAAACGCCGCTGCAAAGTATCTCGTCATCGTGTTC
CTGATCAATGCTCTGGCATGCCGCCCTCGCTTACAAAAAGTTTACTTCATTCTGGTCAACC
TGCTCAAGGCCGCTCTGTTTATCATTTCAATAAAACGCCGAGCTAAGTTGGCCGC
ATGGACCTGTAAGGCCGCTGCAAATTCACTCTCGTGAATCTGCTCATTTTACAACCTCAA
GACGAGGAAAATATCGGAATTATAAGCTGCCCTACGGGAGGACAAACCTGAAAGCCGCTGC
AGTCCTGCTCGGCCGGAGTGGGCTGGTGTCAATTCTGATCTCTTGTATCTGTCCTGGTG
AAGGCCGTCCTGCCGCCCTCGCGAGTCGTGTGA (SEQ ID NO:118)

FIGURE 18H

PfCTL_3 (SEQ ID NO:119)

MQVQIQSLFLLLLWVPGSRGVFLIFFDLFLNAAAPSDGKCNLYKAAAVTCNGIQVRKLFHIFDGD
NEIKAHVLSHNSYEKNEYGKQENWYSLKKILSVFFLANAAAKFIKSLFHIFKAAALYISFYFIKAKF
VAAWTLKAAAKAAAYIIPHQSSLKAAAGLIMVLSFL

ATGGGAATGCAGGTGCAGATCCAGAGCCTGTTCTGCTCCTCTGGGTGCCGGATCCAGA
GGAGTGTCTTGATCTCTTGACCTGTCAGCTGACCTGTGGAAACGGGATTCAAGTCAGGAACACTCTTCAC
AATCTCTACAAGGCCGCTGCAGTGACCTGTGGAAACGGGATTCAAGTCAGGAACACTCTTCAC
ATCTCGACGGCGATAACGAGATCAAGGCCATGTGCTGTCCACAATTCTATGAAAAAAAC
TACTATGGAAGCAAGAGAATTGGTACAGCCTGAAGAAAATTCTGTCCTGTTCTTCGCCCC
AACGCCGCTGCAAAGTTATCAAGTCTGTCCATATTCAAGGCCGCTGCACTCACATCA
GCTTCTATTATTAAAGCCAATTGTGGCCGCTTGGACACTGAAGGCCGCTGCAAAAGCCG
CTGCATACTATATCCCTACCAGAGCTCCCTGAAGGCCGCTGCAGGGCTGATCATGGTGTCT
CTTCCTGTGA (SEQ ID NO:120)

PfCTL/HTL(N) (SEQ ID NO:121)

MQVQIQSLFLLLLWVPGSRSSVFNVVNSSIGLIMVLSFLGPGPGLYISFYFILVNLLIFHINGKIIKN
SEGPGPGPDSIQDSLKESRKLSGPGPGLVAGLLGVVSTVLLGGVGLVLGPGPGLPSENERGYYIPHQ
SSLGPGPGQTNFKSLLRNLGVSENIFLKGPGPQFQDEENIGIYGPGPQKYLVIVFLIFFDLFLVGPGP
GKFIFKSLFHIFDGDNIEGPGPQKSKYKLATSVLAGLLGPGPGLPYGKTNLGPGPGRHNWVNHA
AMKLIGPGPGMRKLAILSVSSFLFVEALFQEYGPGPVTCNGIQVRGPQGMNYYGKQENWYSL
KKGPGPQPSDGKCNLYADSAWENVKNVIGPFMKAVCVEVGPGPKILSVFFLALFFIIFNKGPQPG
HVLSHNSYEKGPGPQKYLQIAGGIAGGLALLACAGLAYKFVVPGAATPYAGEPAPF

ATGGGAATGCAGGTGCAGATCCAGAGCCTGTTCTGCTCCTCTGGGTGCCGGATCCAGA
GGAAGTAGTGTGTTCAATGTTGTGAACTCATCAATTGGTCTGATCATGGTCTGAGCTTCTCG
GCCAGGGCCAGGATTATATATTCTTCTACTCATCCTGTCAACCTGTAATATTCCACAT
TAACGGAAAATAATAAAGAACAGTGAAGGCCCTGGCCCTGGGCTGACTCGATCCAGGATT
CTCTAAAAGAACATCGAGGAAGCTCCGGACCAGGCCCTGGTGTACTGCCGGGTTGCTGGGA
GTAGTTAGCACAGTGTGTTAGGAGGCCTGGCCCTCGCTTAGGACCTGGACCAGGTCTGCCG
TCCAAAACGAAAAGAGGAACTACATACCTCACCAGAGCAGCCTCGGCCAGGCCGGACA
AACCAATTCAAATCCCTTGCAGAACATCTAGGAGTGAGCGAGAACATATTCTAAAGGACC
CGGTCCCGGCTTCAGGACGAGGAGAACATAGGTATTACGGTCCAGGACCTGGAAAATACCT
AGTATCGTATTCTAATTGACCTATTCTGGGGCCCAGGTCCGGAAAGTTCATT
AAATCACTCTTCCACATTGACGGAGATAACGAGATAGGACCCGGTCCGGAAATCAA
GTACAAACTAGCCACTTCAGTGTGGCCGCTTCTAGGGCCGGCCAGGGCTCCCTATGG
AAAGACAAATCTGGCCCGGTCCAGGACGGCACAACCTGGGTGAATCATGGGTTCCATTGG
CCATGAAACTAATCGGGCCCGGTCCAGGACGGCACAACCTGGGTGAATCATGGGTTCCATTGG
TTCTGTTCTAGAGGCCTGTTCAAGAATATGGCCAGGACCTGGGTACATGTGGGAATG
GGATCCAGGTGAGAGGACCGGGACCTGGTATGAACTATTACGGTAAACAGGAAAATTGGTAC
TCCCTGAAAAGGGTCCAGGCCCGGCCCTCAGATGGTAAGTCAACCTGTATGCTGACTCA
GCATGGGAGAACGTAAAAATGTAATAGGCCATTCAAGGGCAGTTGTGTCGAAGTCGG
ACCAGGCCAGGAAAATACCTTCTGTTCTCAGCTCTTCTCATCTTCAACAAG
GGACCAGGGCCAGGTACCGTGTATCCATAACTCTTATGAAAAGGGCCAGGACCTGGGA
ATACAAAATCGCAGGAGGGATCGCCGGCGGGTAGCGCTCCTGCCTGCGCAGGCTGGCTTA
CAAATCGTTGACCAGGAGCTGCAACACCCTATGCAGGAGAACCTGCCATTGAAAGATC
TGC (SEQ ID NO:122)

FIGURE 18I

Pf33 (SEQ ID NO:123)

MGMQVQIQSLFLLLLWVPGSRGMKAVCEVNVTCGNGIQVRKGJIMVLSFLNAALFHIFDGN
EIKAALLACAGLAYKKSFLFVEALFNAAPSDGKCNLYKAAQTNFKSLLRNLPSENERGYKAAGVS
ENIFLKNAAYFILVNLLIKAAAIALSVSSFLFVNTPYAGEPAPFKAAAKYKLATSVLKAAVFLIFFDL
FLNYYIPHQSSLKAAGLLGVNSTVGAVLGGVGLVNLACAGLAYKKAKFIKSLFHIFKAAFYFIL
VNLLKAFLIFFDLFLVKALFFIIFNKNYYGKQENWYSLKFVEALFQEYNAAAKFVAAWTLKAAAK
ILSVFFLANAVLAGLLGVNFQDEENIGIYKAAALYISFYFIKAFILVNLLIFHNAALPYGRTNLKAA
HVLSHNSYEKNAAKYLIVVF

GCCGCCACCATGGGAATGCAGGTGCAGATCCAGAGCCTGTTCTGCTCCTCTGTGGGTGCC
GGATCCAGAGGATTATGAAAAGCTGTCTGTAGAGGTGAATGTAACATGCGGTAACGGAAT
TCAGGTGAGAAAGGGACTCATCATGGTACTCAGCTTCTGAACGCAGCCCTGTCACATCTT
TGACGGAGACAATGAAATCAAAGCCGCATTGCTGCCCTGCCCCACTAGCCTATAAAAAGA
GTTCTTCTGTTGAAGCACTATTAAACGCAGCACCCAGTGACGGTAAATGCAACCTATATA
AAGCAGCTCAGACTAATTCAAAAGCCTGTTAAGAAATCTGCCCTCAGAGAATGAAAGGGT
TACAAAGCCGCCGGCGTGTCCGAGAATATTCTGAAAGAACGCCGTGCTTATTATACTC
GTGAATCTACTCATAAAGGCAGCCGAATCCTTCAGTGTCCAGCCTCTGTTGTAACACAC
CATATGCGGGCGAGCCGGCTCTTCAAGGCTGCAGCAAATACAAGCTGCCACATCAGTAT
TGAAAGCAGCTGTGTTTGATATTCTTGTACTTTAAACTACTACATACCTCATCAGTCT
AGTCTAAAGCAGCCGGCTACTGGGAACGCTCTACTGTGGGGGCCGTCTACTGGAGGA
GTGGCCTCGTGTGAACCTCGCGCAGGTCTGGCTACAAAAAAAGCGAAATTCACTCAAG
TCTCTGTTCCACATTAAAGCCGCATTCTATTCTACTAGTGAACCTCTCAAAGCTTCT
GATCTCTCGATCTATTCTCGTAAAGCGCTATTCTCATTATCTTAAACAAAAATTATTAC
GGCAAGCAAGAAAATTGGTACTCACTCAAGTTGTAGAAGCTGTCCAGGAATACAAGCC
GCTGCTAAATTGCGTGCAGCTGGACCTGAAAGCAGCTGCAAAGATCTATGGTCTCTTC
TCGCTAATGCCGTATTAGCAGGACTCTAGGCAACGTGAACCTTCAAGACGAAGAGAATATAG
GCATCTACAAAGCCGCAGCACTGTACATTCTACTTCATCAAGGCCCTCATACTGGTCAA
CCTTCTGATATTCTACATATGCAGCACTGCCATATGGGAGAACCAACTTGAAGCGGCCACGT
GTGAGGCCACAACCTCTACAGAGAAGAACGCCGCCGAAATATCTGTCATTGTCCTGAT
TTGA (SEQ ID NO:124)

TB.1 (SEQ ID NO:125)

MMQVQIQSLFLLLWVPGSRRGRMSRVTTFTVKALVLLMLPVVNLIGTAAAVVKALVLLMLPVGA
GLMTAVYLVGAAAMALLRLPVKRMFAANLGVNLSYFGGICVGRLPLVLPVNAAAAKFVAAWT
LKAAGAKAARLMIGTAAAGFVVALIPLVNAMTYAAPLFVGAAAAMALLRLPLV

ATGCAGGTGCAGATCCAGAGCCTGTTCTGCTCCCTGTGGGTGCCCGGATCCAGAGGAAGG
ATGAGCAGAGTGACCACATTCACTGTCAAGGCCCTGGTCTCCTGATGCTCCCCGTGTAAC
CTGATGATCGGCACCGCTGCAGCCGTGAAAGCTCTCGTCCTGCTCATGCTCCCTGTGGGA
GCAGGGCTGATGACAGCCGTACCTGGTGGCGCTGCAGCCAATGGCCCTCCTGCGGGCTGCCA
GTGAAGCGCATGTTGCTGCAAATCTGGAGTCAACTCCCTATTTGGGGCATTTGCGTG
GGAAGGCTGCCCTCGTGCCTGCTGTGAATGCAGCCGTGCCAAATTGTCGCCGCTTGG
ACTCTGAAGGCAGCCGCTAAGGCCGCTGCAAGACTGATGATCGGGACC CGCTGCCGGCTT
CGTGGTCGCCCTGATTCCCTGGTGAACGCCATGACATACGCAGCTCCTGTTGTGGGAGC
CGCTGCAGCCATGGCTCCTGCGGCTGCCACTGGTGTGA (SEQ ID NO:126)

FIGURE 18J

BCL A2 #90 (SEQ ID NO:127)

MQVQIQSLFLLLWVPGSRGIMIGHLVGVNRLQETELVNAKVAEIVHFLNAKVFGSLAFVNAYL
SGANLNVGAAYLQLVFGIEVNAAAKFVAAWTLKAAAKAAAVVLGVVFGINSMPPPGRVNAAA
ATVGIMIGVNAKLCPVQLWV

ATGCAGGTGCAGATCCAGAGCCTGTTCTGCTCCTCTGTGGGTGCCCGGGTCCAGAGGAATT
ATGATCGGCCATCTGGTGGCGTCACAGACTGCTGCAGGAAACCGAGCTGGTAATGCCAA
GGTGGCCGAAATTGTGCACCTTCTCAACGCAAAGGTGTTGGTCCCTGGCTTTGTCAATGCC
TATCTGAGCGGCGCTAACCTCAACGTCGGAGCCGCTACCTCCAGCTGGCTTCGGCATCGAG
GTCAACGCTGCTGCAAATTCTGTCAGCTGGACCTCAAGGCTGCAGCAAAGGCTGCCGCC
GTCGTGCTCGGAGTGGTGTCCGGATCAACTCTATGCCACCTCCCGGACTAGGGTCAATGCT
GCCGCCGCAACAGTGGGAATCATGATTGGGTGAATGCCAAACTGTGCCAGTGCAACTGTG
GGTGTGA (SEQ ID NO:128)

BCL A2 #88 (SEQ ID NO:129)

MQVQIQSLFLLLWVPGSRGVVLGVVFGINAAAAKFVAAWTLKAAAKVAEIVHFLNAYLSGANL
NVGAAYLQLVFGIEVNIMIGHLVGVNRLQETELVNAKVFGLAFVNNAKLCPVQLWVNAAAATV
GIMIGVNSMPPPGRV

ATGCAGGTGCAGATCCAGAGCCTGTTCTGCTCCTCTGTGGGTGCCCGGGTCCAGAGGAGTC
GTGCTGGGAGTCGTCTCGGCATTAATGCCCGCTGCAAAGTCGTGGCTGCCCTGACCCCTG
AAGGCCGCACTAAAGTGGCAGAGATCGCACCTTCTGAACGCCCTACCTGAGCGGAGCAA
TCTGAACGTCGGCGCTGCCTATCTGCAGCTGTGTTGGATTGAAGTGAACATCATGATTGG
ACATCTGGTGGCGTGAACAGGCTGCTCCAGGAAACTGAGCTGGTCAACGCTAAAGTGTTCG
GGTCTCTGCCCTTGTGAACGCTAACGCTCTGCCCGTCAACTCTGGTCAATGCCAGCCG
CTACAGTGGGGATCATGATCGGCGTGAACCTCCATGCCCTACCAGGGACCAGAGTGTGA
(SEQ ID NO:130)

BCL A2 #63 (SEQ ID NO:131)

MQVQIQSLFLLLWVPGSRGKLCPVQLWVNAAAATVGIMIGVNIIGHLVGVNRLQETELVNA
KVAEIVHFLNAKVFGSLAFVNAYLSGANLNVGAAYLQLVFGIEVNAAAKFVAAWTLKAAAKAA
AVVLGVVFGINSMPPPGRV

ATGCAGGTGCAGATCCAGAGCCTGTTCTGCTCCTCTGTGGGTGCCCGGGTCCAGAGGAAAG
CTCTGCCCGTGCAACTGTGGGTCAACGCCGCCGCCAACCGTCGGCATTATGATCGGGGTG
AACATCATGATCGACACCTGGTCGGCGTGAACAGGCTGCTGCAGGAGACAGAACTGGTCAA
TGCCAAGGTGGCTGAAATTGTCCATTCTGAATGCCAAAGTGTTCGGCTCTCGCTTCGTG
AACGCTTATCTGAGCGGAGCTAACCTCAACGTGGGGGCCGATACCTCCAGCTCGTCTTGGG
ATTGAGGTGAATGCCGAGCTAAATTGTGCGTGCCTGGACCTGAAGGCAGCAGCCAAGGCT
GCCGCAGTGGTGTGGAGTGGTGTGGATTCAATTCCATGCCCTACCAGGCAGTAGAGTG
TGAGGATCC (SEQ ID NO:132)

FIGURE 18K

Prostate 1 (SEQ ID NO:133)

LTFFWLDRSVKAAAVLVPQWVLTVKAAALLQERGVAYIKAALLSIALSVNPLVCNGVLQGVK
AAIMYSAHDTTVKAAFLPKLQCVNAMMNDQLMFLNAGLPSIPVHPVKAALGTTCYVGAAI
LLWQPIPVNFLRPRSLQCVKAFLTSVTWIGVNALLYSLVHNLAATLMSAMTNL

ATGCAGGTGCAGATCCAGAGCCTGTTCTGCTCCCTGTGGGTGCCCGGGTCCAGAGGGATTG
ACATTTTTGGCTGGATAGATCGGTAAGGCTGCAGCCGTGCTTGTTCATCCCCAGTGGGTCT
TGACCGTAAAGGCTGCCGCGTCTACAAGAAAGAGGGTGCATACTCAAAGCTGCTCTC
CTCTTGAGTATTGCGCTAAGTGTAAACCCGCTAGTTGTAATGGGGTGTACAAGGTGTGAAA
GCGGCAGATTATGTACAGTCCCCACGACACTACCGTAAAAGCAGCCGCTTCTGACCCCAAA
AAACTCCAATGCGTAACCGAATGATGAATGATCAGCTGATGTTAAACGCTGGCTTACCT
TCTATACCGGTTCATCCAGTCAGGCGCGGCAATTGGGTACGACGTGTTATGGAGCAGCG
ATACTCTTGGCAGCCCATAACCAAGTAAATTAAAGACCTAGATCCTTACAATGCGTCAAAG
CATCCCTAACACTCTAGTAACCTGGATCGGAGTCAATGCTCTGCTATATAGCCTCGTACACAA
CTTGGCGCGGCCACACTTATGAGTGCAATGACGAATTAGCTAAGTCGTGGCGGCTGGAC
TCTAAAGGCCGCAGCA (SEQ ID NO:134)

HIV-1043 (SEQ ID NO:135)

MEKVYLAWVPAHKGIGGGPGPGQKQITKIQNFRVYYRGPGPGWEFVNTPLVKLWYQGPQGYR
KILRQRKIDRLIDGPQPGQHLLQLTVWGIKQLQGPQPGGEIYKRWIILGLNKIVRMYGPQGQQM
VHQAIISPTLNQPGPGIKQFINMWQEVGKAMYGPQPGWAGIKQEFGIPYNPQGPQPGKTAQVMA
VFIHNFKRGPGPGSPAIFQSSMTKILEPGPGPGEVNIVTDSQYALGIQGPQGHNSWRAMASDFNLPP
GPGGAETFYVDGAANRETKGPGPGGAVVQDNSDIKVVPGPQPGFRKYTAFTIPSINNE

ATGGAGAAGGTGTACCTGGCTGGTTCCAGCCCCACAAAGGCATCGGGGGAGGGCCGGACC
TGGCAGAAACAGATCACCAAGATCAGAACTTCCGGGTATACTACCGGGGACCTGGTCCAG
GTTGGAGTTGTGAACACACCACCTTAGTAAAGCTCTGGTACCAAGGGCCCCGGTCCGGAT
ACCGTAAAATCCTGAGGCAAAGAAAGATAGATCGCCTCATGGATGGCCCAGGCCAG
CACCTCTGCAGCTTACAGTGTGGGAATTAAACAGCTGCAGGGGCCGGCCCCGGGGGG
AATTATAAAAGGTGGATCATTCTGGGTCTGAACAAGATCGTCCGCATGTATGGCCCTGGACC
CGGACAGGGCAGATGGTCACCAAGCAATCAGCCCTCGAACCTGAATGGACCGGGCCAG
GAATCAAGCAATTCAATTAAACATGTGGCAAGAAGTTGGTAAGGCTATGTACGGTCCCGGCCCTG
GATGGCAGGGATAAAACAGGAGTTGAATCCCTACAATCCCAGGGTCTGGGCCAGGT
AAAACGGCAGTGCAGATGGCGTGTCAATTATAAGCGGGGCCCTGGACCTGGCAGC
CCAGCTATATTCAAAGTTCGATGACCAAAATCTGGAGCCCAGGGCCGGCGAAGT
GAACATTGTACAGATTCTAGTATGCCCTCGCATAGGGCCGGACCAGGGCATTCAA
TTGGCGGCCATGGCGTCTGACTTAATCTACCTCTGGCCAGGGCCTGGCGCGAAACTT
CTATGTGGACGGCGCTGCAAACAGGGAGACTAAGGGACCCGGACCCGGCGCTGTAGTCA
TTCAGGACAACACTAGACATCAAGGTGGTCCCGGTCCAGGGCCGGITCAGAAAGTATACCG
CCTTCACTATTCCGTCCATCAACAATGAGTGA (SEQ ID NO:136)

FIGURE 18L

HIV-1043 PADRE (SEQ ID NO:137)

MEKVYLAWVPAHKIGGGPGPGQKQITKIQNFRVYRGPGPGWEFVNTPPLVKLWYQGPGPGYR
KILRQRKIDRLIDGPQPGQHQLLQLTVWGIKQLQGPQGEYKRWIILGLNKIVRMYGPQPGQGQM
VHQAISPRTLNGPGPGIKQFINMWQEVGKAMYGPQPGWAGIKQEFQIPYNPQGPQPGKTAQVMA
VFIHNFKRGPGPGSPAIFQSSMTKILEPGPGPGEVNIVTDSQYALGIIGPGPGHSNWRAMASDFNLPP
GPGPGGAETFYVDGAANRETKGPGPGGAVVIQDNSDIKVVPGPQGPFRKYTAFTIPSINNEGPGPGA
KFVAATLKA

ATGGAGAAGGTGTACCTGGCCTGGGTTCCAGCCCACAAAGGCATGGGGGGAGGGCCCGGACC
TGGGCAGAACAGATCACCAAGATCCAGAACACTTCGGGTATACTACCGGGGACCTGGTCCAG
GTTGGGAGTTGTGAACACACCACCTTAGTAAAGCTGGTACCAGGGCCCCGGTCCCAGGAT
ACCGTAAAATCTGAGGCAGAACAGAAAGATAGATCGCCTCATGATGGCCCCGGCCAGGCCAG
CACCTTCTGCAGCTTACAGTGTGGGAATTAAACAGCTGCAGGGGCCGGCCCCGGGGGGA
AATITATAAAAGGTGGATCATTCTGGGTCTGAACAAGATCGTCCGATGTATGGCCCTGGACC
CGGACAGGGGCAGATGGTCCACCAAGCAATCAGCCCTCGAACCTTGAATGGACCGGGCCAG
GAATCAAGCAATTCTTAACATGTGGCAAGAACAGTTGTAAGGCTATGTACGGTCCCCGGCCTG
GATGGGCAGGGATAAAACAGGAGTTGGAATCCCTAACATCCCCAGGGTCTGGCCAGGT
AAAACGGCAGTCAGATGGCGTGTTCATTCTATAATTAAAGCGGGGCCCTGGACCTGGCAGC
CCAGCTATATTCAAAGTTCGATGACCAAAATCTGGAGGCCGCCAGGGCCGGCGAAGT
GAACATTTGTACAGATTCTCAGTATGCCCTCGGCATCATAGGGCCGGACCAGGGCATTCAA
TTGGCGGCCATGGCGTCTGACTTTAATCTACCTCTGGCCAGGCCCTGGCGCGAAACTTT
CTATGTGGACGGCGCTGCAAACAGGGAGACTAAGGGACCCGGACCCGGCGCTGTAGTCA
TTCAGGACAACCTAGACATCAAGGTGGTCCAGGCCCGGGTTAGAAAGTATACCG
CCTTCACTATTCCGTCCATCAACAATGAGGGCCCCGGCCAGGTGCCAAGTTCGTGGCTGCCT
GGACCCCTGAAGGCTGCCGCTTGA (SEQ ID NO:138)

HIV 75mer (SEQ ID NO:139)

EKVYLAWVPAHKIGGPGQGQMVKQISPRTLNGPGSPAIFQSSMTKILEPGPGPGFRKYTA
FTIPSINNE

GAGAAGGTGTACCTGGCCTGGGTGCCACAAGGAATCGGAGGACCTGGCCCTGGACA
GGGACAGATGGTGCACCAGGCCATAGCCCTAGGACCTGAACGGACCTGGACCTGAAAGCC
CTGCCATCTTCAGAGCAGCATGACCAAGATCCTGGAGCCGGACCTGGACCTGGATTAGGA
AGTACACCGCCTCACCATCCCCAGCATCAACAACGAGTGA (SEQ ID NO:140)

FIGURE 18M

PfHTL (SEQ ID NO:141)

MQVQIQSLFLLLLWVPGSGRHNVNHA VPLAMKLIGPGPGKCNLYADSAWENVKNGPGPGKS
KYKLATSVLAGLLPGPGQTNFKSLRNLGVSEGPGPGSSVFNVVNSSIGLIMGPGPVKNVIGPF
MKAVCVEGPGPGMNYYGKQENWYSLKKGPGPGLAYKFVVPGAATPYGPGPGPDSIQDSLKESR
KLNGPGPGPLLIFHINGKIIKNSEGPGAGLLGNVSTVLLGGVGPGPKYKIAGGIAGGLALLGPGP
GMRKLAISVSSFLV

ATGGGAATGCAGGTGCAGATCCAGAGCCTTTCTGCTCCTCCTGTGGGTGCCGGATCCAGA
GGAAGGCACAACCTGGGTGAATCATGCTGTGCCCTGGCTATGAAGCTGATCGGCCCTGGACC
AGGGAAATGCAACCTCTACGCAGACAGCCTGGAGAACGTCAAGAATGGCCCCGGACCTG
GGAAATCCAAGTATAAGCTCGTACCTCTGTGCTGGCAGGCCTGCTGGACCAGGCCCGGAC
AGACAAAATTCAAAGCCTGCTCAGAAACCTGGAGTGTCGAGGGCCCTGGCCCAGGATCT
AGCGTCTTAATGTGGTCAACTCCTCTATTGGGCTCATCATGGGACCCGGACCTGGGTGAAA
AATGTCATTGGCCCATTCATGAAGGCCGTGTGTGCGAAGGACCCGGCCTGGCATGAAC
TATGGAAAGCAAGAAAATTGGTACAGCCTGAAGAAAGGCCCTGGCCAGGCAGACTGGCTTA
CAAGTTGTGGTCCCAGGGCAGCCACTCCCTATGGGCTGGCCAGGCCGATTCCATCCA
GGACTCTCTCAAAGAGAGCCGGAAACTGAACGGACCCGGCCTGGACTGCTCATTTCCACAT
CAATGGCAAATTATCAAGAACAGCAGGGACCTGGCCAGGCAGGACTGCTGGGAACG
TGTCCACCGTCTGCTGGCGAGTGGGCCCTGGGAAGTACAAGATCGCTGGAGGG
ATCGCAGGCGGACTGGCCCTCTGGGCCAGGACCAGGGATGCGCAAACGGCTATTCTCT
GTCTCCAGTTCTGTTGTGA (SEQ ID NO:142)

FIGURE 18N

Protein	Sequence(SEQ ID NOS:143-340)	Restriction
HIV gag 386	VLAEAMSQV	HLA-A2
HIV gag 271	MTNNPPIPV	HLA-A2
HIV pol 774	MASDFNLPPV	HLA-A2
HIV pol 448	KLVGKLNWA	HLA-A2
HIV pol 163	LVGPTPVNI	HLA-A2
HIV pol 498	ILKEPVHGV	HLA-A2
HIV pol 879	KAACWWAGI	HLA-A2
HIV pol 132	KMIGGIGGFI	HLA-A2
HIV pol 772	RAMASDFNL	HLA-A2
HIV pol 183	TLNFPISPI	HLA-A2
HIV env 134	KLTPLCVTL	HLA-A2
HIV env 651	LLQLTVWGI	HLA-A2
HIV env 163	SLLNATDIAV	HLA-A2
HIV nef 221	LTFGWCFL	HLA-A2
HIV vpr 59	AIRILQQL	HLA-A2
HIV vpr 62	RILQQLLFI	HLA-A2
HIV pol 929	QMAVFHNFK	HLA-A3
HIV pol 722	KVYLAWVPAHK	HLA-A3
HIV pol 971	KIQNFRVYYR	HLA-A3
HIV pol 347	AIFQSSMTK	HLA-A3
HIV pol 98	VTIKIGGQLK	HLA-A3
HIV env 61	TTLFCASDAK	HLA-A3
HIV env 47	VTVYYGVPVWK	HLA-A3
HIV nef 100	QVPLRPMTYK	HLA-A3
HIV vif 7	VMIVWQVDR	HLA-A3
HIV gag 162	QMVKHQAISSPR	HLA-A3
HIV gag 545	YPLASLRSLF	HLA-B7
HIV gag 237	HPVHAGPLA	HLA-B7
HIV pol 186	FPISPIETV	HLA-B7
HIV pol 893	IPYNPQSQGVV	HLA-B7
HIV env 259	IPIHYCAPA	HLA-B7
HIV env 250	CPKVSFEPI	HLA-B7
HIV nef 94	FPVRPQVPL	HLA-B7
HIV rev 75	VPLQLPPL	HLA-B7
HIV pol 684	EVNIVTDSQY	HLA-A1
HIV gag 317	FRDYVDRFY	HLA-A1
HIV pol 368	VIYQYMDDLY	HLA-A1
HIV pol 295	VTVLVDVGDAY	HLA-A1
HIV pol 533	IYQEPMFKNL	HLA-A24
HIV pol 244	PYNTPVFAI	HLA-A24
HIV pol 530	TYQIYQEPP	HLA-A24
HIV pol 597	YWQATWIPEW	HLA-A24
HIV env 681	IWGCSGKLI	HLA-A24
HIV env 671	RYLKDDQLL	HLA-A24

FIGURE 19A

Protein	Sequence	Restriction
HIV env 55	VWKEATTLF	HLA-A24
HIV vpr 46	IYETYGDTW	HLA-A24
HIV vpr 14	PYNEWTLEL	HLA-A24
HIV gag 298	KRWIILGLNKIVRMY	HLA-DR
HIV pol 596	WEFVNTPPLVKLWYQ	HLA-DR
HIV pol 956	QKQITKIQNFRVYYR	HLA-DR
HIV pol 712	KVYLAWVPAHKGIGG	HLA-DR
HIV gag 294	GEIYKRWIILGLNKI	HLA-DR
HIV pol 711	EKVYLAWVPAHKGIG	HLA-DR
HIV env 729	QHLLQLTVWGIKQLQ	HLA-DR
HIV gag 171	QGQMVFHQAIISPTLN	HLA-DR
HIV pol 335	SPAIFQSSMTKILEP	HLA-DR
HIV env 566	IKQFINMWQEVGKAMY	HLA-DR
HIV pol 303	FRKYTAFTIPSINNE	HLA-DR
HIV pol 758	HSNWRAMASDFNLPP	HLA-DR
HIV pol 915	KTAVQMAVFIHNFKR	HLA-DR
HIV vpu 31	YRKILRQRKIDRLID	HLA-DR3
HIV pol 874	WAGIKQEFGIPYNPQ	HLA-DR3
HIV pol 674	EVNIVTDSQYALGII	HLA-DR3
HIV pol 619	AETFYVDGAANRETK	HLA-DR3
HIV pol 989	GAVVIQDNSDIKVVP	HLA-DR3
HCV NS4 1812	LLFNILGGWV	HLA-A2
HCV NS1/E2 728	FLLLADARV	HLA-A2
HCV NS4 1590	YLVAYQATV	HLA-A2
HCV NSS 2611	RLLVFPDLGV	HLA-A2
HCV CORE 132	DLMGYIPLV	HLA-A2
HCV NS4 1920	WMNRLIAFA	HLA-A2
HCV NS4 1666	VLVGGVLAA	HLA-A2
HCV NS4 1769	HMWNFISGI	HLA-A2
HCV NS4-1851	ILAGYGAGV	HLA-A2
HCV CORE 35	YLLPRRGPRL	HLA-A2
HCV NS1/E2 726	LFLLLLADA	HLA-A2
HCV LORF 1131	YLVTRHADV	HLA-A2
HCV CORE 51	KTSERSQPR	HLA-A3
HCV CORE 43	RLGVRATRK	HLA-A3
HCV ENV1 290	QLFTFSPRR	HLA-A3
HCV NS1/E2 632	RMYVGGVEHR	HLA-A3
HCV NS3 1396	LIFCHSKKK	HLA-A3
HCV NS4 1863	GVAGALVAFK	HLA-A3
HCV NS4 1864	VAGALVAFK	HLA-A3
HCV NS3 1262	LGFGAYMSK	HLA-A3
HCV Core 169	LPGCSFSIF	HLA-B7
HCV NSS 2922	LSAFSLHSY	HLA-A1
HCV NS3 1128	CTCGSSDLY	HLA-A1
HCV NSS 2180	LTDPSSHITA	HLA-A1

FIGURE 19B

Protein	Sequence	Restriction
HCV Core 126	LTCGFADLMGY	HLA-A1
HCV NS3 1305	LADGGCSEGGAY	HLA-A1
HCV NS4 1765	FWAKHMWNF	HLA-A24
HCV NS5 2875	RMILMTHFF	HLA-A24
HCV NS5 2639	VMGSSYGF	HLA-A24
HCV NS4 1765	FWAKHMWNFI	HLA-A24
P. falciparum SSP2-230	FMKAVCVEV	HLA-A2
P. falciparum EXP1-83	GLLGVVSTV	HLA-A2
P. falciparum CSP-7	ILSVSSFLFV	HLA-A2
P. falciparum LSA1-94	QTNFKSLLR	HLA-A3
P. falciparum LSA1-105	GVSENIFLK	HLA-A3
P. falciparum SSP2-522	LLACAGLAYK	HLA-A3
P. falciparum SSP2-539	TPYAGEPAPF	HLA-B7
P. falciparum LSA1-1663	LPSENERGY	HLA-A1
P. falciparum EXP1-73	KYKLATSVL	HLA-A24
P. falciparum CSP-12	SFLFVEALF	HLA-A24
P. falciparum LSA1-10	YFILVNLLI	HLA-A24
P. falciparum SSP2-14	FLIFFDLFLV	HLA-A2
P. falciparum EXP1-80	VLAGLLGVV	HLA-A2
P. falciparum EXP1-91	VLLGGVGLVL	HLA-A2
P. falciparum SSP2-523	LACAGLAYK	HLA-A3
P. falciparum EXP1-10	ALFFIIFNK	HLA-A3
P. falciparum LSA1-11	FILVNLLIFH	HLA-A3
P. falciparum SSP2-126	LPYGRTNL	HLA-B7
P. falciparum CSP-15	FVEALFQEY	HLA-A1
P. falciparum LSA1-1794	FQDEENIGIY	HLA-A1
P. falciparum LSA1-9	FYFILVNLL	HLA-A24
P. falciparum SSP2-8	KYLVIVFLI	HLA-A24
P. falciparum CSP-394	GLIMVLSFL	HLA-A2
P. falciparum EXP1-2	KILSVFFLA	HLA-A2
P. falciparum CSP-344	VTCGNGIQVR	HLA-A3
P. falciparum LSA1-59	HVLSHNSYEK	HLA-A3
P. falciparum SSP2-207	PSDGKCNLY	HLA-A1
P. falciparum LSA1-1671	YYIPHQSSL	HLA-A24
P. falciparum LSA1-1876	KFIKSLFHIF	HLA-A24
P. falciparum SSP2-13	VFLIFFDLFL	HLA-A24
P. falciparum LSA1-1881	LFHIFDGDNEI	HLA-A24
P. falciparum CSP-55	YYGKQENWYSL	HLA-A24
P. falciparum LSA1-5	LYISFYFI	HLA-A24
P. falciparum CSP-2	MRKLAILSVSSFLV	HLA-DR
P. falciparum CSP-53	MNYYGKQENWYSLKK	HLA-DR
P. falciparum CSP-375	SSVFNVVNSSIGLIM	HLA-DR
P. falciparum SSP2-61	RHNWVNHAVPLAMKLI	HLA-DR
P. falciparum SSP2-165	PDSIQDSLKESRKLN	HLA-DR3
P. falciparum SSP2-211	KCNLYADSAWENVKN	HLA-DR3

FIGURE 19C

Protein	Sequence	Restriction
P. falciparum SSP2-223	VKNVIGPFMKAVCVE	HLA-DR
P. falciparum SSP2-509	KYKIAGGIAGGLALL	HLA-DR
P. falciparum SSP2-527	GLAYKFVVPGAAATPY	HLA-DR
P. falciparum EXP1-71	KSKYKLATSVLAGLL	HLA-DR
P. falciparum EXP1-82	AGLLGNVSTVLLGGV	HLA-DR
P. falciparum LSA1-16	LLIFHINGKIIKNSE	HLA-DR
P. falciparum LSA1-94	QTNFKSLLRNLLGVSE	HLA-DR
HBV core 18	FLPSDFFPSV	HLA-A2
HBV env 183	FLLTRILTI	HLA-A2
HBV env 335	WLSLLVPFV	HLA-A2
HBV pol 455	GLSRYVVARL	HLA-A2
HBV pol 538	YMDDVVLGV	HLA-A2/A1
HBV pol 773	ILRGTSFVYV	HLA-A2
HBV pol 562	FLLSLGIHL	HLA-A2
HBV pol 642	ALMPLYACI	HLA-A2
HBV env 338	GLSPTVWLSV	HLA-A2
HBV core 141	STLPETTVVRR	HLA-A3
HBV pol 149	HTLWKAGILYK	HLA-A3/A1
HBV pol 150	TLWKAGILYK	HLA-A3
HBV pol 388	LVVDFSQFSR	HLA-A3
HBV pol 47	NVSIPWTHK	HLA-A3
HBV pol 531	SAICSVVRR	HLA-A3
HBV pol 629	KVGNFTGLY	HLA-A3/A1
HBV pol 665	QAFTFSPTYK	HLA-A3
HBV core 19	LPSDFFPSV	HLA-B7
HBV env 313	IPIPSSWAF	HLA-B7
HBV pol 354	TPARVTGGVF	HLA-B7
TB	RMSRVTTFV	HLA-A2
TB	ALVLLMLPVV	HLA-A2
TB	LMIGTAAAVV	HLA-A2
TB	ALVLLMLPV	HLA-A2
TB	GLMTAVYLV	HLA-A2
TB	MALLRLPV	HLA-A2
TB	RMFAANLGV	HLA-A2
TB	SLYFGGICV	HLA-A2
TB	RLPLVLPAV	HLA-A2
TB	RLMIGTAAA	HLA-A2
TB	FVVALIPLV	HLA-A2
TB	MTYAAPLFV	HLA-A2
TB	AMALLRLPV	HLA-A2
p53 139	KLCPVQLWV	HLA-A2
CEA 687	ATVGIMIGV	HLA-A2
CEA 691	IMIGHLVGV	HLA-A2
Her2/neu 689	RLLQETELV	HLA-A2
MAGE3 112	KVAEIVHFL	HLA-A2

FIGURE 19D

Protein	Sequence	Restriction
Her2/neu 369	KVFGSLAFV	HLA-A2
CEA 605	YLSGANLNV	HLA-A2
MAGE2 157	YLQLVFGIEV	HLA-A2
Her2/neu 665	VVLGVVFVGI	HLA-A2
p53 149	SMPPPGTRV	HLA-A2
PAP.21.T2	LTFFWLDERSV	HLA-A2
PAP.112	TLMSAMTNL	HLA-A2
PAP.284	IMYSAHDTTV	HLA-A2
PSM.288.V10	GLPSIPVHPV	HLA-A2
PSM.441	LLQERGVAYI	HLA-A2
PSM.469L2	LLYSLVHNL	HLA-A2
PSM.663	MMNDQLMFL	HLA-A2
PSA.3.V11	FLTLSVTWIGV	HLA-A2
PSA.143.V8	ALGTTCYV	HLA-A2
PSA.161	FLTPKKLQCV	HLA-A2
HuK2.4.L2	LLLSIALSV	HLA-A2
HuK2.53.V11	VLVHPQWWLTV	HLA-A2
HuK2.165	FLRPRSLQCV	HLA-A2
HuK2.216.V11	PLVCNGVLQGV	HLA-A2

FIGURE 19E